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Thermoplastic hybrid-matrix composites prepared by a room-temperature vacuum infusion and in-situ polymerisation process

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Abstract

This work explores a novel route for the fabrication of hybrid-matrix composites based on a recently developed liquid thermoplastic acrylic resin. This liquid resin was modified using a poly(phenylene ether) (PPE) oligomer with vinyl functionality. Glass fibre-reinforced laminates based on acrylic and PPE-modified acrylic matrices were produced by a room-temperature vacuum infusion and in-situ polymerisation process. Comparative assessments of their mechanical performance and mode-I interlaminar fracture behaviour revealed enhanced matrix ductility, transverse flexural properties and initiation fracture toughness. Crazing was identified as the dominant mechanism for improved resistance to crack initiation.

1 Introduction

Innovative low-viscosity liquid thermoplastic (LTP) resins can readily infiltrate into fibrous reinforcement under conditions of relatively low temperature and pressure in the same way that thermoset (TS) resins can [1–4]. Room-temperature infusible acrylic resins with viscosities as low as 100 mPa.s have received considerable research attention in recent years [5–13]. In our previous work, we presented comparisons between the mechanical performance of acrylic composites with equivalent epoxy composites and reported inferior transverse flexural performance [7] and impact damage resistance [8] in the acrylic-matrix composites.

Structural composites typically comprise a thermoset matrix or a semi-crystalline thermoplastic matrix. Cross-linked networks and crystalline domains contribute to enhanced matrix rigidity, making them ideal candidates for high-performance applications. In contrast, purely amorphous matrices such as acrylics do not contain cross-links or crystalline regions within their molecular structure. Thus, this might influence composite properties, particularly when matrix strength plays a key role.

Therefore, there is a significant scope to tailor the structure of acrylic-matrix composites for enhanced performance under different loading conditions. Recent works on this topic have used Nanostrength™ triblock copolymers comprising polymethylmethacrylate-b-polybutylacrylate-b-polymethylmethacrylate [9–11] and hybrid fibre reinforcements [12] to realise improved composite properties. However, TP-TP hybridisation of an acrylic matrix, via in-situ polymerisation, is novel and never investigated before.

Poly(phenylene ether) (PPE) – an amorphous engineering thermoplastic, is arguably one of the most successfully applied as a modifier in TS-matrix composites [14–16]. Unlike the acrylic matrix, which is a purely aliphatic amorphous TP, PPE contains aromatic rings, which may confer some rigidity in a hybrid system and is thus, worthy of exploration.

This present study investigates an innovative route to obtaining vacuum-infusible hybrid-matrix composites based on acrylic and PPE. To promote reactive blending during in-situ polymerisation of the hybrid matrix, PPE with vinyl functionality was selected for this study. The effects of hybridisation on mechanical and morphological properties are presented herein.

2 Experimental

2.1 Materials and fabrication

Two 4-mm thick (nominally) test laminates were prepared by a room-temperature vacuum infusion and in-situ polymerisation process. Table 1 provides an overview of the materials used. Full details of the materials and the fabrication processes used are supplied in Appendix A.

Table 1. Summary of materials used for composite fabrication.

	<i>Elum[®] 188 O^a</i>	<i>NORYL[™] SA9000^b</i>	<i>Q-UD Glass^c</i>
<i>Unreinforced polymer samples^d</i>			
<i>A100/P0</i>	100	0	0
<i>A95/P5</i>	95	5	0
<i>Composite samples^d</i>			
<i>GF/A100/P0</i>	100	0	50
<i>GF/A95/P5</i>	95	5	57

^a A Liquid acrylic resin [A] supplied by Arkema GRL, France.
^b An oligomeric PPE resin [P] with vinyl functionality, supplied by SABIC.
^c TEST2594 – a quasi-unidirectional (UD) glass non-crimp fabric (NCF) supplied by Ahlstrom-Munksjö. GF: glass fibre. Fibre volume fraction.
^d Polymerised using a dibenzoyl peroxide initiator – BP50FT supplied by United Initiators.

2.2 Mechanical and thermomechanical characterisation

2.2.1 Tensile testing

Tensile properties were evaluated in accordance with ASTM D3039 under transverse tension.

2.2.2 Short beam shear testing

Short beam shear properties were evaluated by short beam shear testing using a span-to-thickness ratio of 4:1 in accordance with ASTM D2344.

2.2.3 Flexural testing

Non-standard flexural testing was performed on unreinforced matrix samples as detailed in Appendix B. To gain further insights on differences in fracture behaviour of the matrices, SEM inspections were also performed.

Flexural properties of glass fibre-reinforced composite samples were determined by three-point bending (ASTM D7264 – Procedure A) using a span-to-thickness ratio of 32:1 under longitudinal and transverse loading.

2.2.4 Mode-I interlaminar fracture toughness (ILFT) testing

Mode-I ILFT was evaluated using double cantilever beam tests per ASTM D5528. SEM inspections were conducted on DCB fracture surfaces to assess fracture behaviour.

The interested reader is referred to Appendix B for supplementary specimen and test specifications.

3 Results and discussions

3.1 Flexural test results of unreinforced matrices

PPE modification appears to improve flexural strength and stiffness of the GF/A95/P5 sample as evidenced by the stress-displacement curves in Figure 1(a). Although mid-span deflections were not measured during testing, the observed increase in stiffness may tentatively indicate an increase in modulus. These results are based on single-sample tests and are thus, not conclusive. These results provide interesting insights, however, that are worthy of further investigation.

The micrographs from the regions of interest, diagrammatically shown in Figure 1(b), reveal relatively flat fracture topography for A100/PO (Figure 1(c)), and multi-planar fractures for the A95/P5 matrix (Figure 1(d)), which suggests an interplay of crack deflection and crack penetration mechanisms as detailed in Appendix C [17]. At higher magnifications, the A100/PO matrix appears homogenous (Figure 1(e)); a biphasic morphology comprising discrete domains was observed for the A95/P5 matrix (Figure 1(f)). These domains are likely PPE-rich phase, surrounded by an acrylic-rich phase.

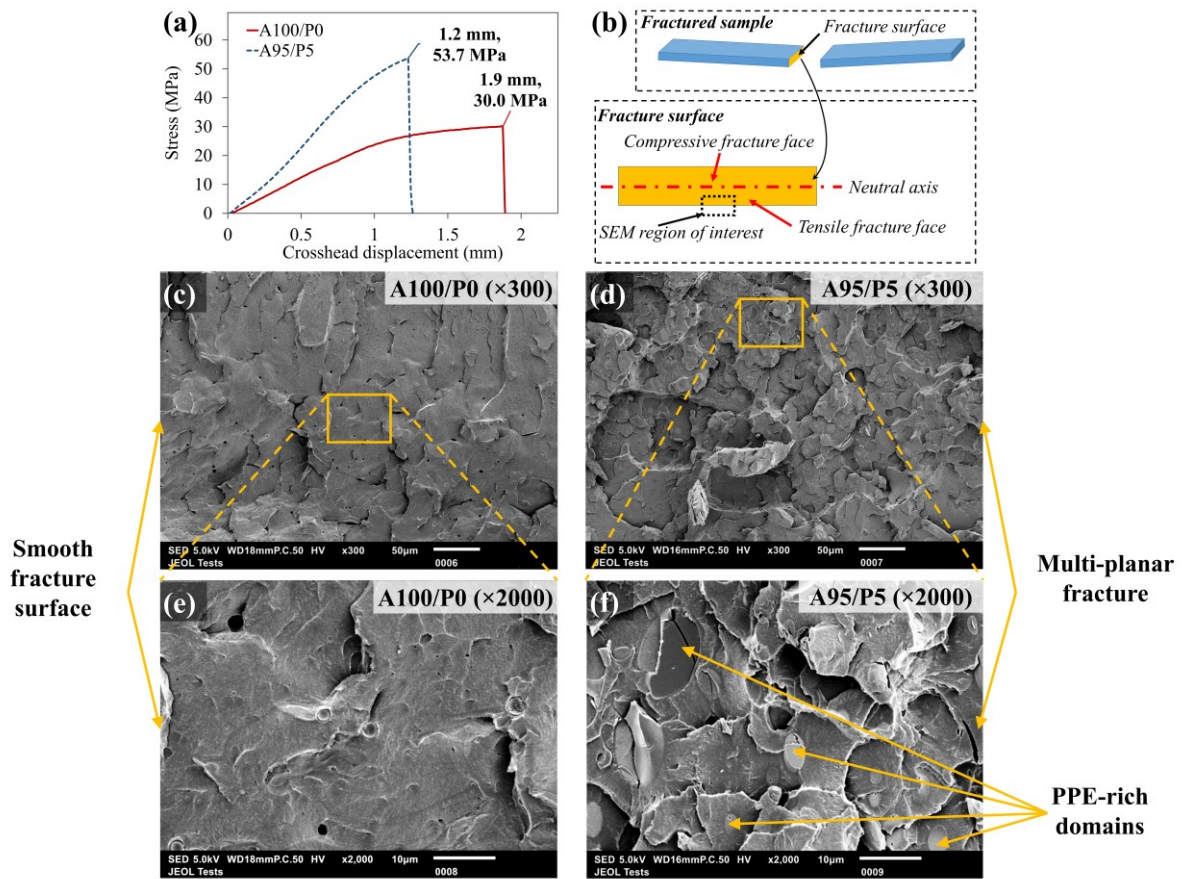


Figure 1. Flexural stress-displacement curves (a) and diagrammatic representation of the SEM region of interest (b). SEM micrographs of fracture surfaces of unreinforced (c) & (e) A100/P0 and (d) & (f) A95/P5 samples at different magnifications ($\times 300$ & $\times 2000$).

3.2 Results of composites testing

3.2.1 Transverse tensile test results

Representative stress-strain responses and average transverse tensile strengths, moduli and failure strains of the GF/A100/P0 and GF/A95/P5 materials are presented in Figure 2(a). Both materials exhibit similar linear behaviour initially; however, an earlier onset of damage initiation (matrix cracking) was observed with the GF/A95/P5 samples. From Points 1 to 3 Figure 2(a), matrix crack accumulation occurs before ultimate failure. In contrast, the GF/A100/P0 material undergoes plastic deformation up to failure. Matrix hybridisation resulted in reduced (-18%) transverse tensile strength with a slight increase in modulus (+8%) and significant increases in failure strain (+58%). Thus, hybridisation appears to increase both transverse composite modulus and ductility. Moreover, higher areas bounded under GF/A95/P5 curves may suggest enhanced toughness.

3.2.2 Short beam shear test results

Figure 2(b) shows the results of short beam shear tests performed on the GF/A100/P0 and GF/A95/P5 materials. For both materials, all samples exhibited plastic deformation up to their respective ultimate shear stress values (Point 1). However, beyond this point, the curves of GF/A95/P5 samples exhibited a more abrupt loss in stiffness with increasing displacement between Points 1 and 2.

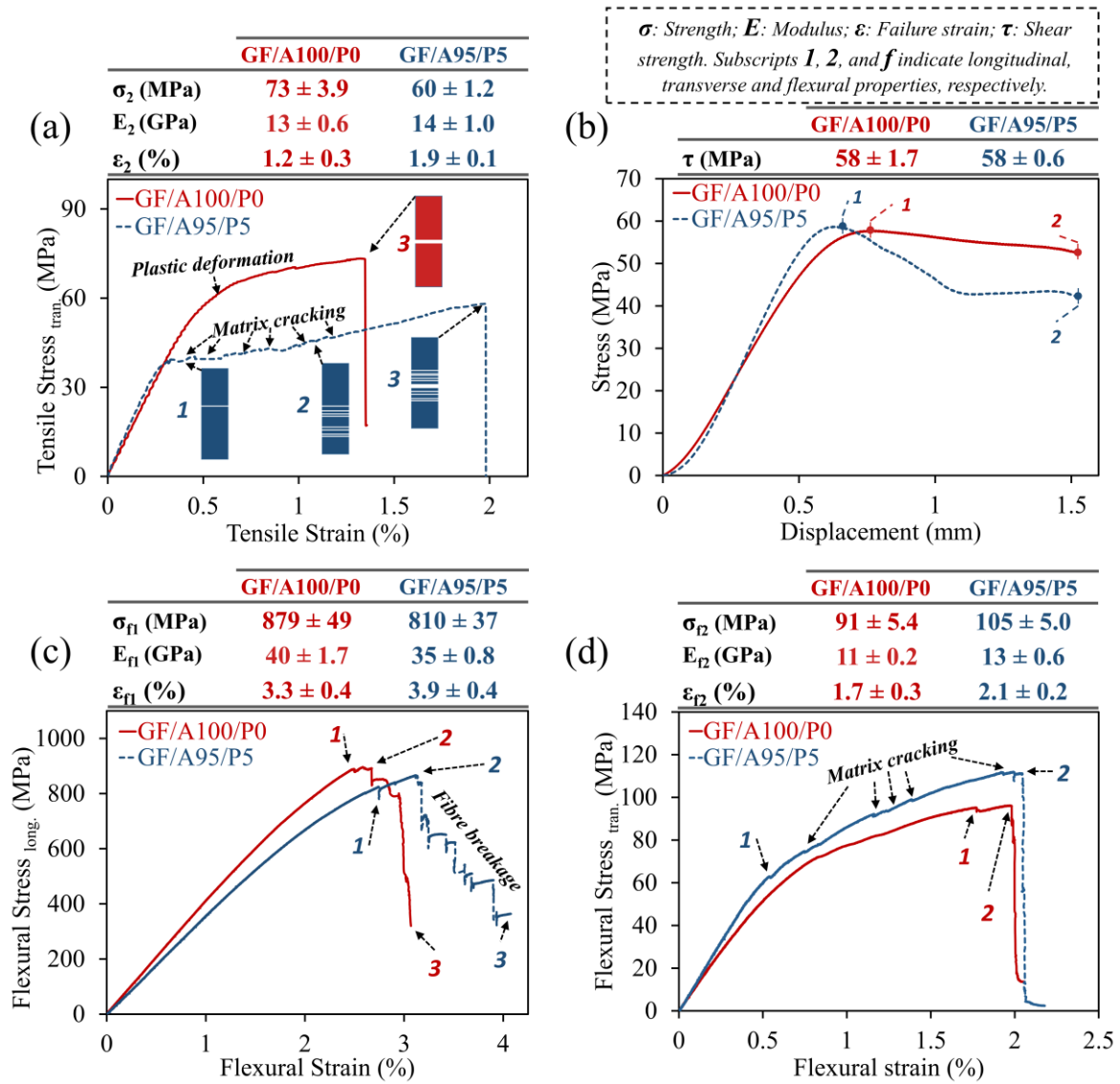


Figure 2. Representative curves and results for GF/A100/P0 (red) and GF/A95/P5 (blue) following loading in (a) transverse tension; (b) short beam shear; (c) longitudinal flexure and (d) transverse flexure.

3.2.3 Flexural test results

Results from longitudinal flexural tests are presented in Figure 2(c). All samples of both materials exhibited a three-stage stress-strain evolution: (i) an initial linear-elastic region, (ii) a region of slight nonlinearity, and (iii) the onset of damage (Point 1). Post-peak strain

evolution between Points 2 and 3 was relatively more confined in GF/A100/P0 samples than in GF/A95/P5. Progressive fibre fractures over a broader range of strains may provide evidence of superior damage resistance and possibly toughness in the GF/A95/P5 material. Moreover, it exhibited markedly higher (18%) average failure strain than the GF/A100/P0.

Hybridisation did, however, produce a laminate with lower longitudinal flexural strength (-8%) and modulus (-18%).

In Figure 2(d), the results of transverse flexural tests are presented. All samples across both materials exhibited an initial region of linearity, beyond which, plastic deformation ensued with a distinct onset of failure (Point 1) and abrupt ultimate failure at Point 2. All GF/A95/P5 samples underwent cumulative matrix cracking in plies under tension, such as those shown between Points 1 and 2. The hybrid-matrix composite exhibited improved transverse flexural strength (+15%), modulus (+18%) and failure strain (+24%) relative to the unmodified reference.

Differences in the trends between the comparative transverse tensile and flexural performance were likely attributed to the sensitivity of the former to defect distribution across the gauge length. Thus, it can be concluded that hybridisation improved the composite transverse strength, modulus, ductility and overall interfacial strength.

3.2.4 Mode-I interlaminar fracture toughness test results

Representative DCB load-displacement curves and obtained results are shown in Figure 3(a). Despite exhibiting superior longitudinal flexural stiffness, GF/A100/P0 samples had unexpectedly lower crack opening stiffness up to initiation, which may be explained by a higher fibre volume fraction in the GF/A95/P5 laminate. Both materials underwent unstable crack growth due to the presence of 90° fibres within the fabric.

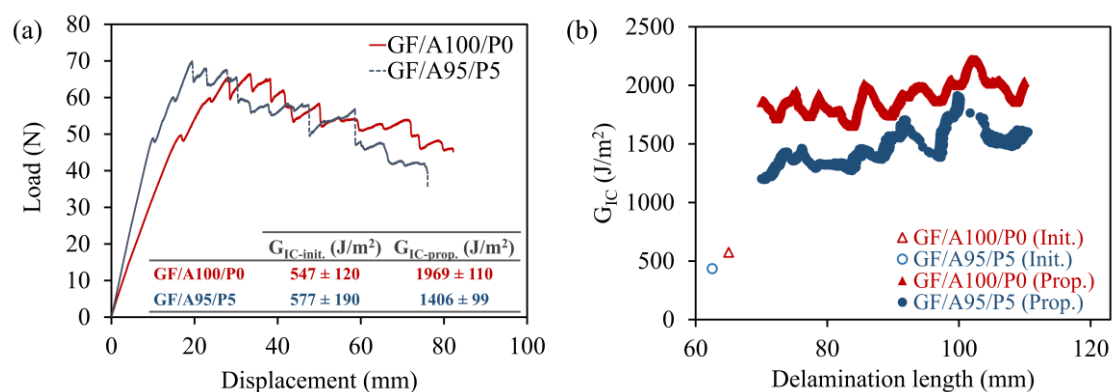


Figure 3. (a) Representative load-displacement curves and (b) R-curves for GF/A100/P0 (red) and GF/A95/P5 (blue) following double cantilever beam testing.

Hybridisation conferred a 5% increase in the initiation fracture toughness ($G_{IC-init.}$); however, propagation fracture toughness ($G_{IC-prop.}$) decreased by 29%. Similar results were reported by Lee et al. [18] who found that hybridisation only enhanced $G_{IC-init.}$, but $G_{IC-prop.}$ was reduced due to limited fibre bridging in the hybrid composite. This is supported by literature on factors affecting propagation behaviour [11,19,20]. Moreover, other factors limiting fibre bridging in the GF/A95/P5 material may be its plausibly higher matrix modulus [17] (evidenced by the higher stiffness reported in 3.1) and enhanced interfacial strength as discussed in 3.2.3 [21,22]. Interestingly, R-curves (Figure 3(b)) did not reveal discernibly distinct propagation behaviour between both materials.

Figure 4 (a)-(f) presents DCB fracture surfaces of GF/A100/P0 and GF/A95/P5 samples obtained using SEM. Both surfaces appear texturally coarse and dull, indicating comparable ductility on a microscopic scale.

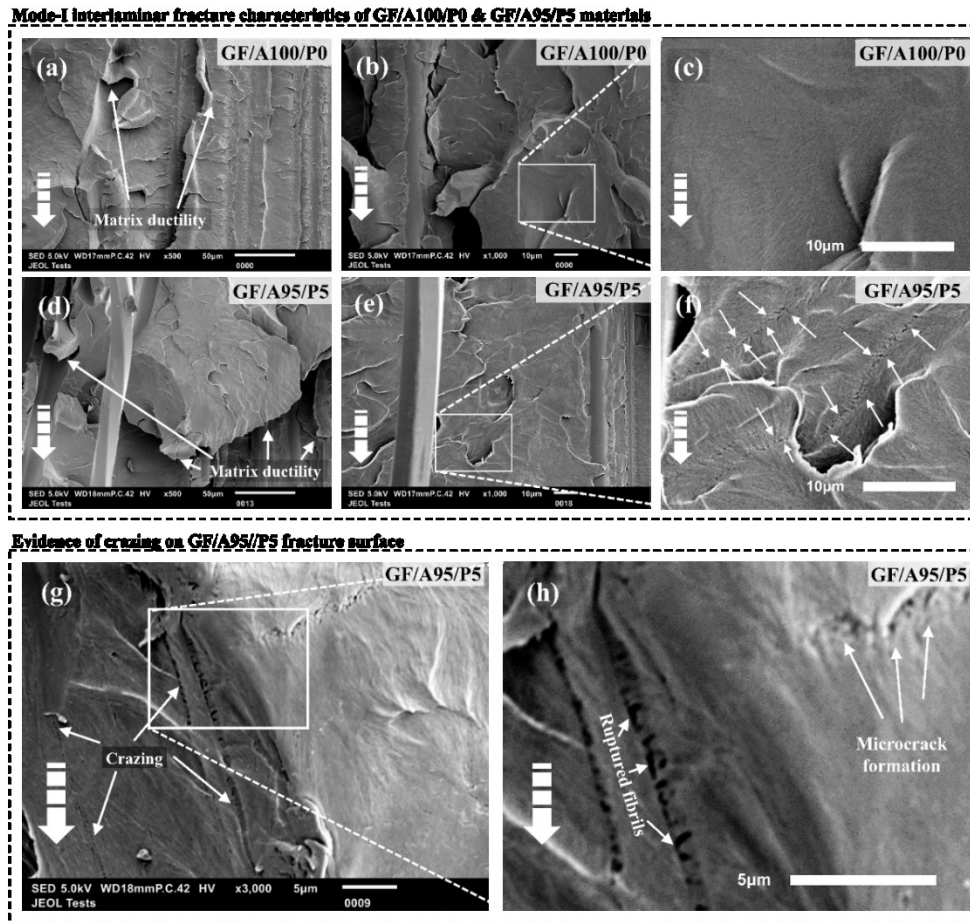


Figure 4. SEM micrographs showing mode-I fracture surfaces of (a), (b) & (c) GF/A100/P0 and (d)-(h) GF/A95/P5. The larger broken arrows show the direction of crack propagation. In (f), arrows highlight paths of microcrack formation.

The GF/A95/P5 sample showed evidence of microcrack formation (Figure 4(f)) and multiple sites of crazing (Figure 4(g)), features which were not observed for GF/A100/Po. The microcracks appeared as long craze-like interpenetrating paths across the fracture surface; however, no coalescence was observed at their points of intersection. Crazing is a dominant plastic deformation mechanism in amorphous TP matrices [23,24], which may explain the increased $G_{IC-init.}$.

The absence of discernible PPE-rich domains in the micrographs of the GF/A95/P5 sample compared with those of the A95/P5 sample may highlight the effects of fibres on the resulting phase morphology. However, further investigations would be required to substantiate this hypothesis.

4 Conclusions

This study represents the first implementation of a novel approach for room temperature vacuum infusion of continuous fibre, thermoplastic hybrid-matrix composites. The approach exploits the low viscosity of liquid TP acrylic resins and with a higher performance poly(phenylene ether) with vinyl functionality to realise enhanced reactivity during the in-situ polymerisation processing. The following are the key observations and conclusions from the benchmarking of mechanical performance with respect to an unmodified acrylic reference laminate:

- **Enhanced ductility in the hybrid-matrix composite:** failure strains increased under transverse tension (+58%), transverse flexure (+24%) and longitudinal flexure (+18%).
- **Improved composite transverse flexural strength (+15%) and modulus (+18%):** this may suggest enhancements in matrix strength, modulus and interfacial adhesion.
- **A 5% increase in initiation fracture toughness,** possibly due to the effects of multiple crazing of the hybrid matrix system.
- **Decreased propagation fracture toughness by 29%,** possibly due to diminished contributions from fibre bridging.

The investigation of the reaction kinetics and mechanism between acrylic resin and PPE, and how this relates to phase separation and morphology is recommended as future work.

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